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#### Bashkirov et al.

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#### (54) METHODS AND APPARATUSES FOR NUCLEIC ACID SHEARING BY SONICATION

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	C12Q 1/68	(2006.01)
	C12P 19/34	(2006.01)
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	C07H 21/04	(2006.01)
	C12N 15/10	(2006.01)

(52) U.S. Cl. CPC .......... *C12Q 1/6806* (2013.01); *C12N 15/1093* 

(58) Field of Classification Search

(2013.01)

See application file for complete search history.

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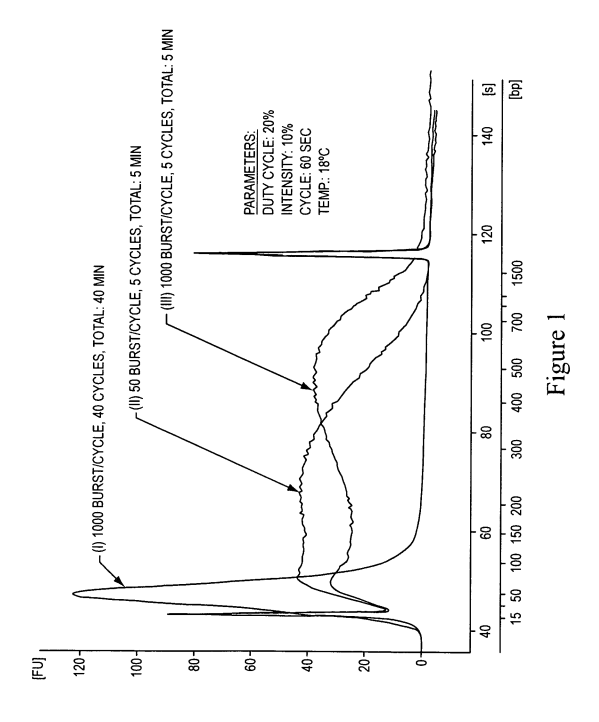
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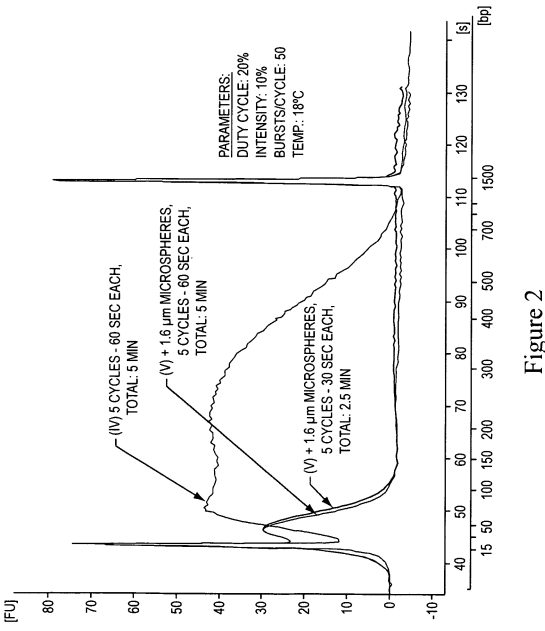
Primary Examiner — Frank Lu

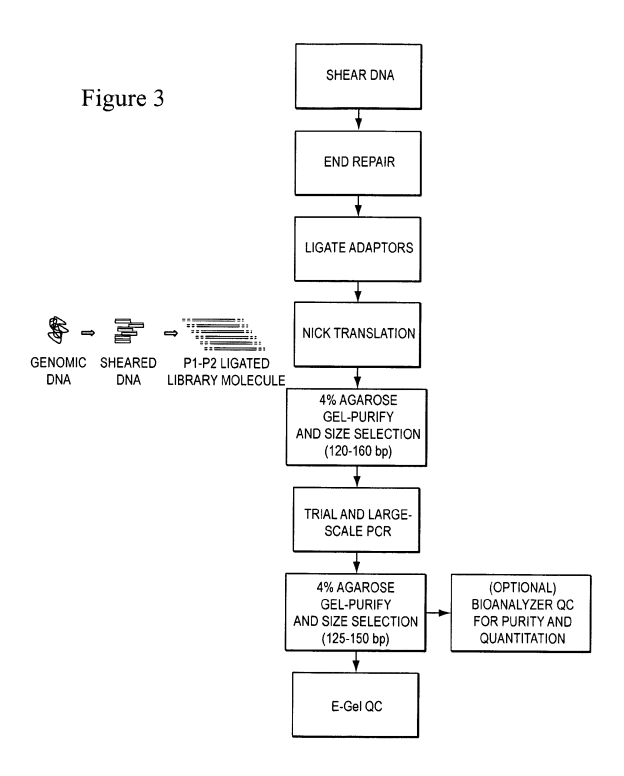
#### (57) ABSTRACT

Methods and kits for preparing nucleic acid fragments from a sample of purified nucleic acid are provided. Alternatively, chromatin or other long polymers can be sheared with similar methods and kits.

#### 15 Claims, 27 Drawing Sheets







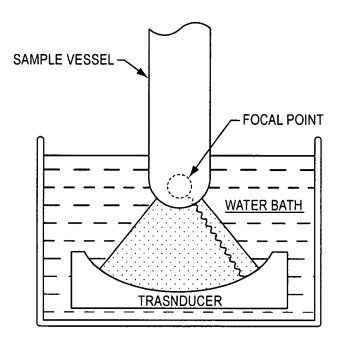


Figure 4

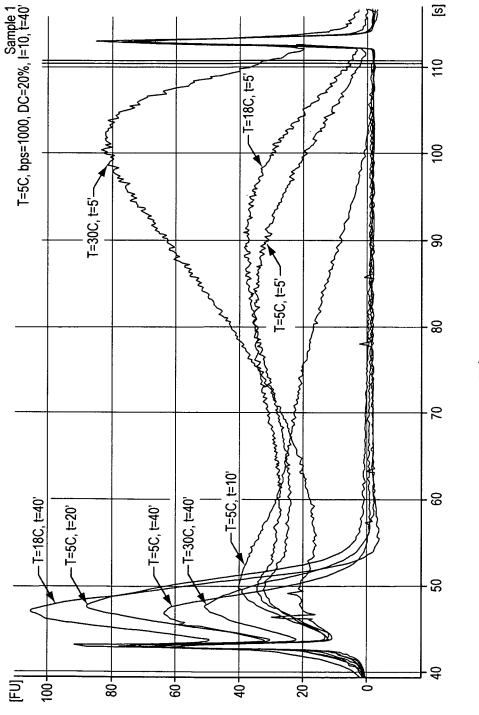


Figure 5

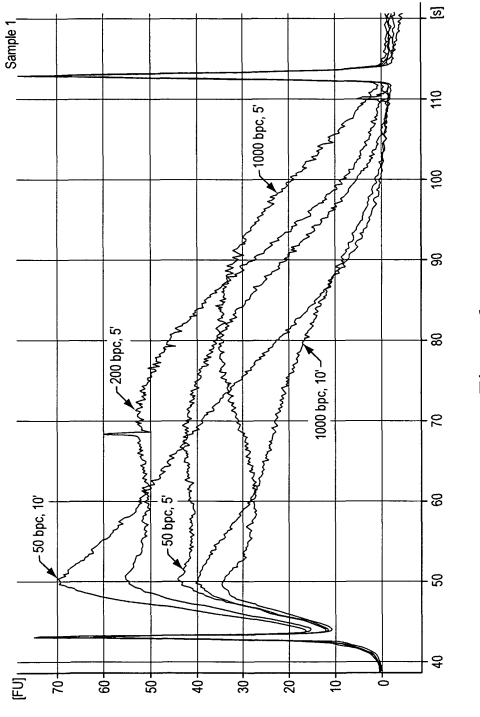
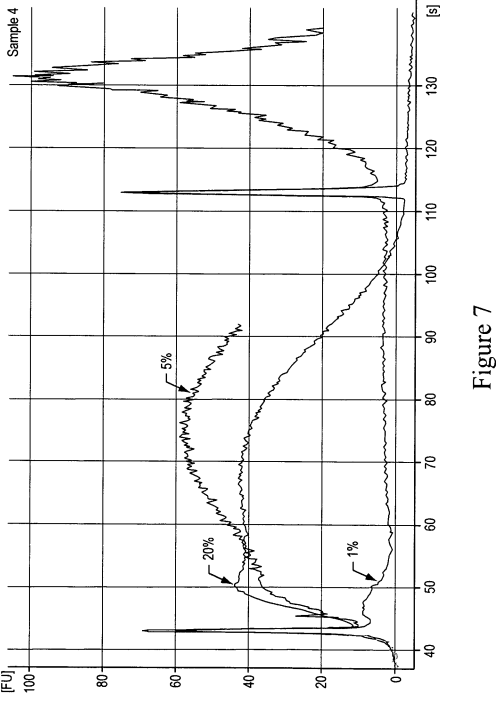
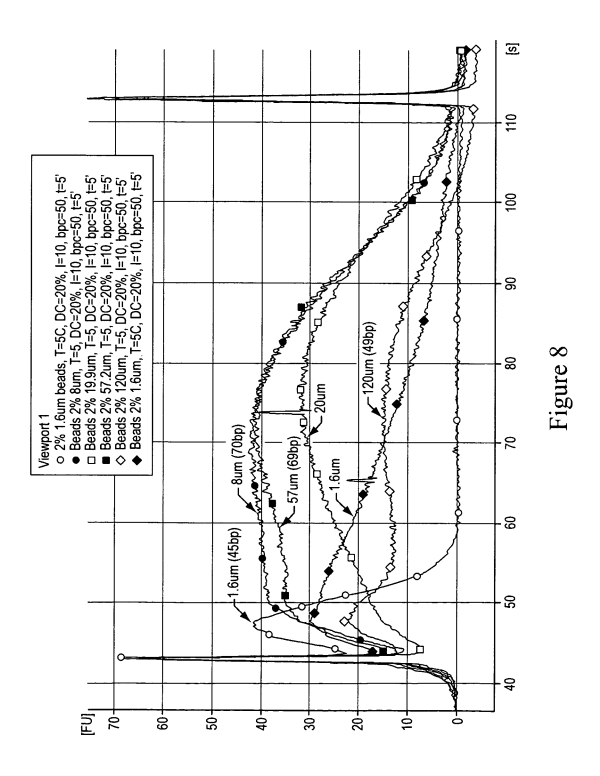
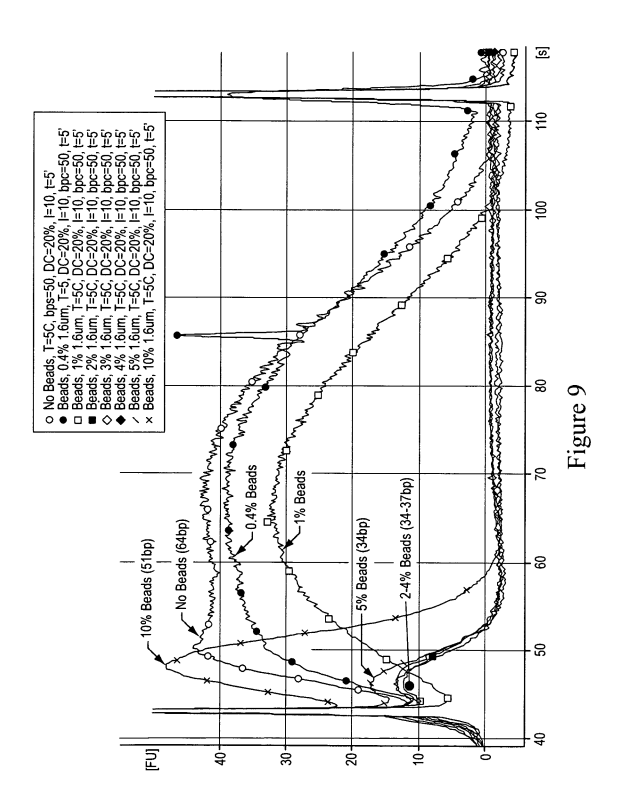
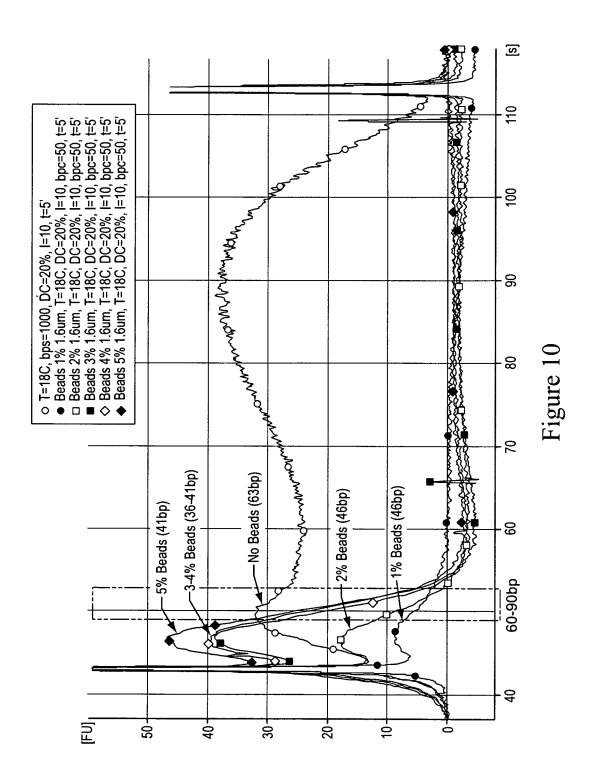


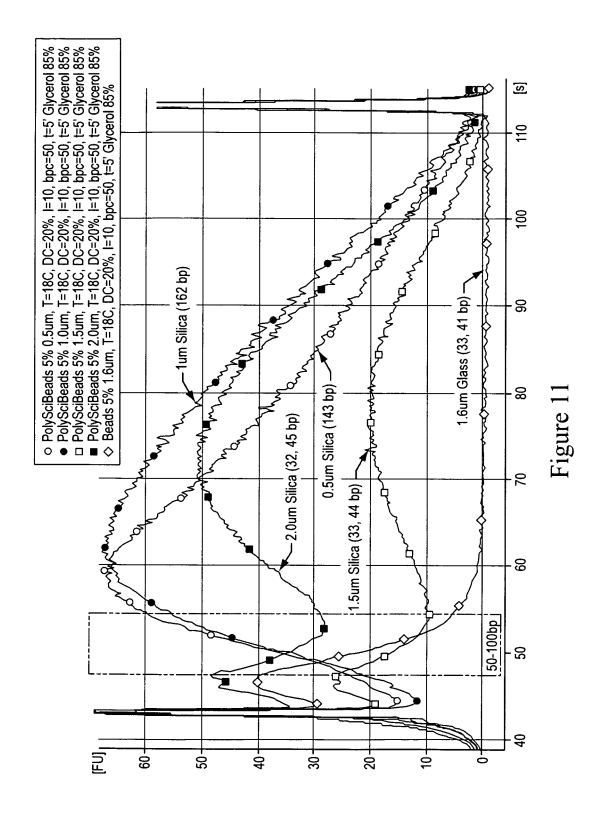
Figure 6











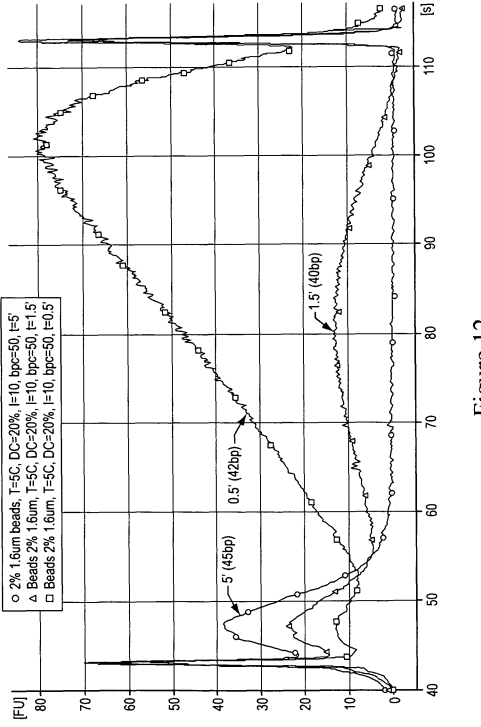
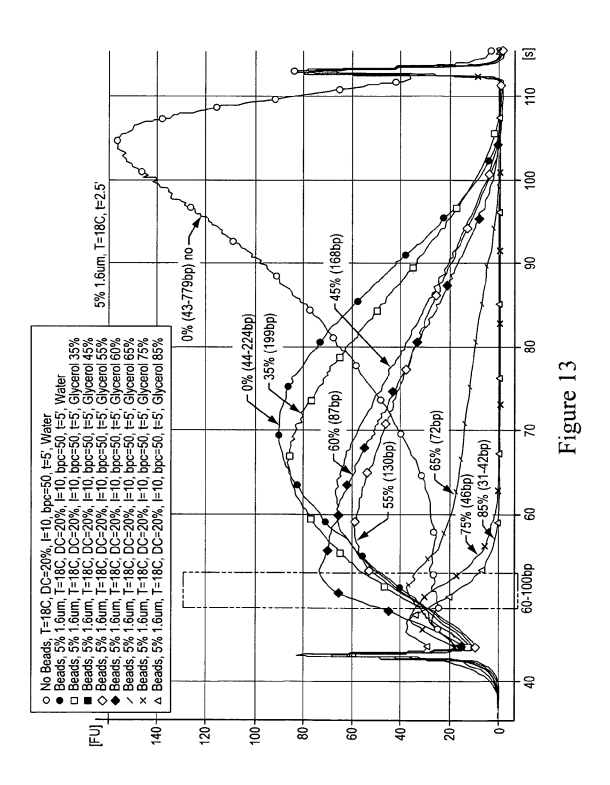
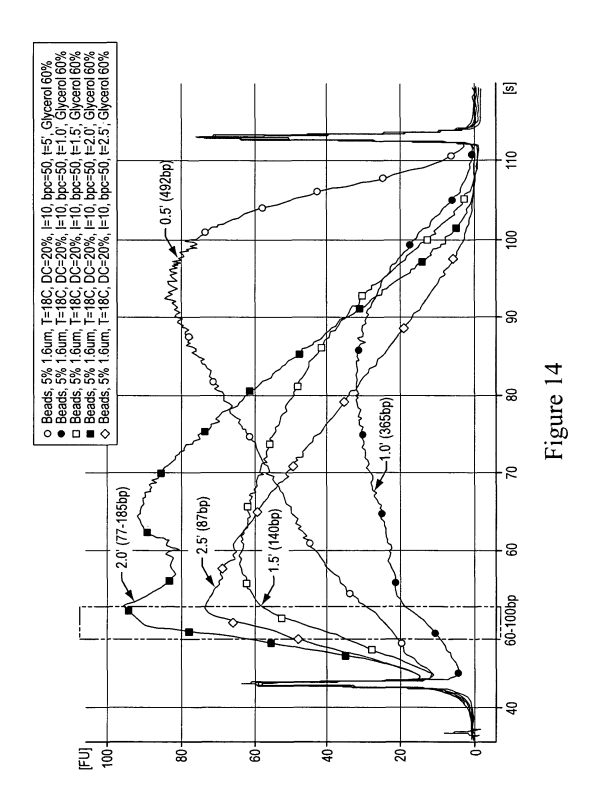


Figure 12





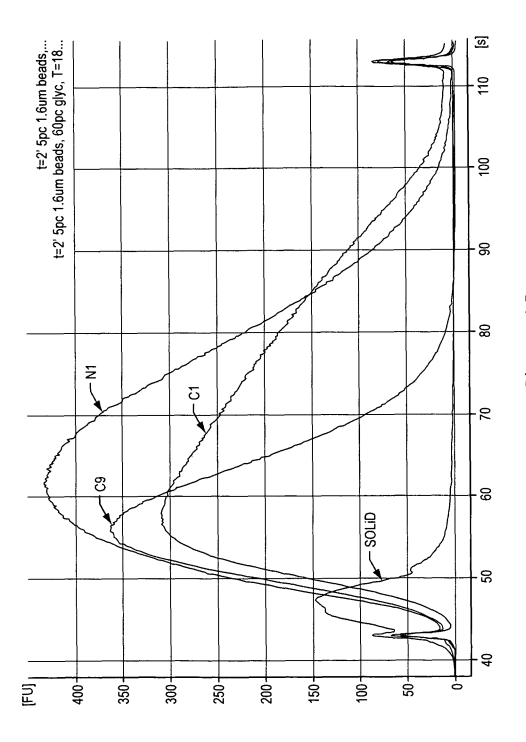


Figure 15

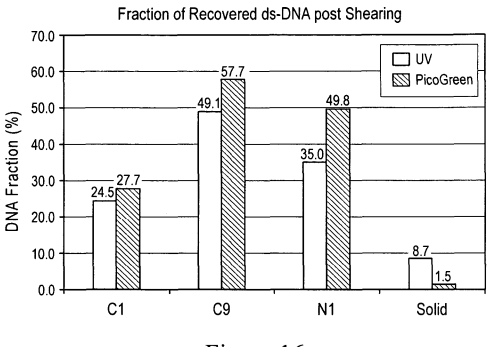
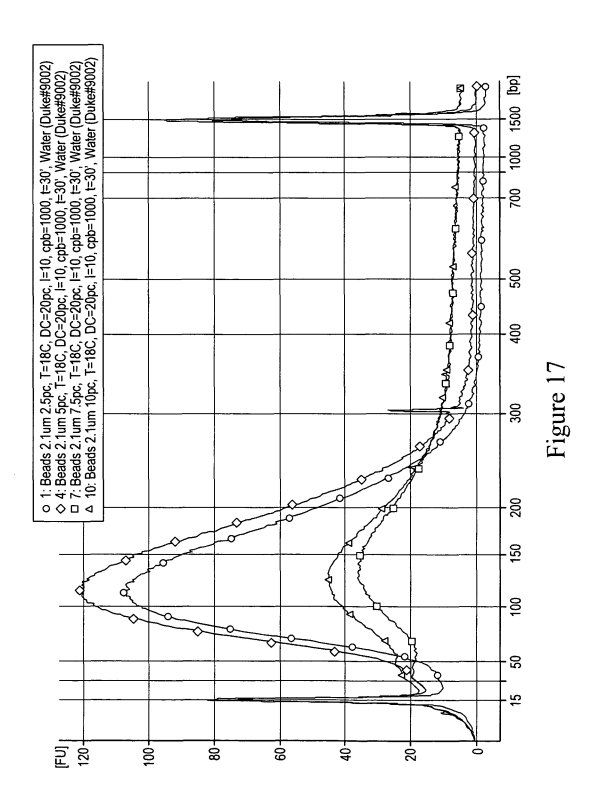
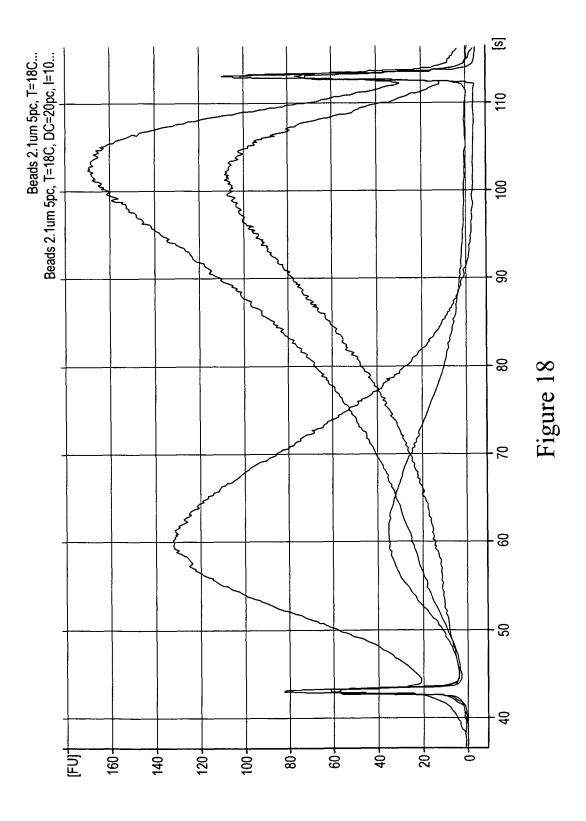
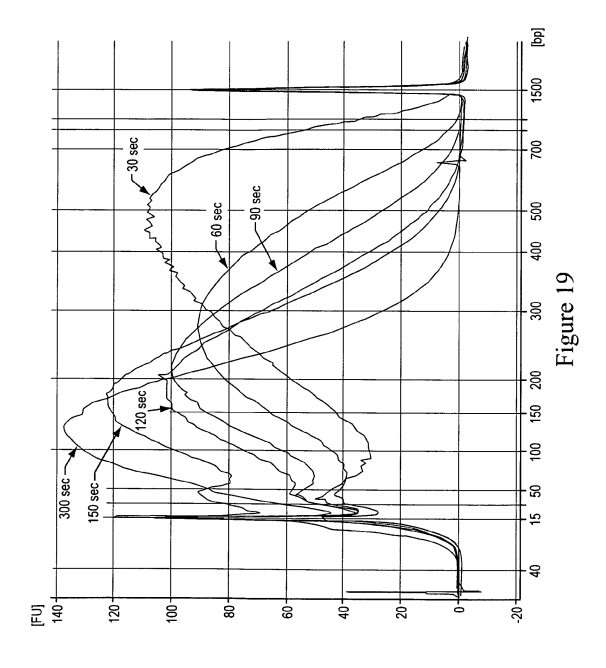
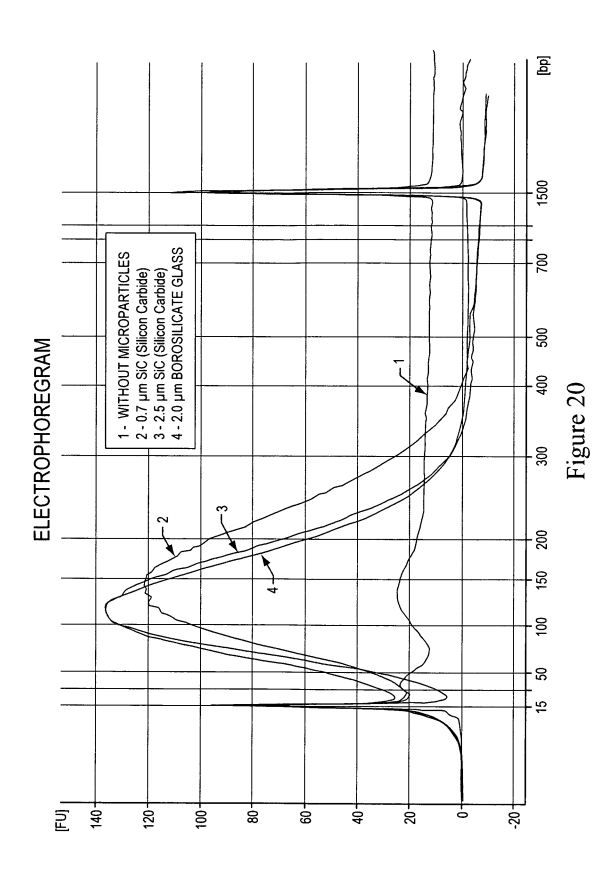


Figure 16









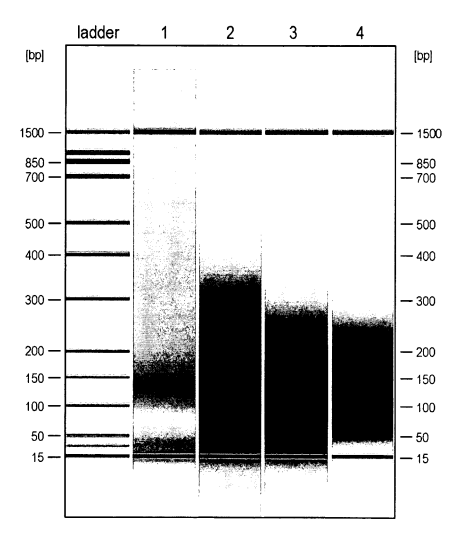


Figure 21

# Matching Summary Report

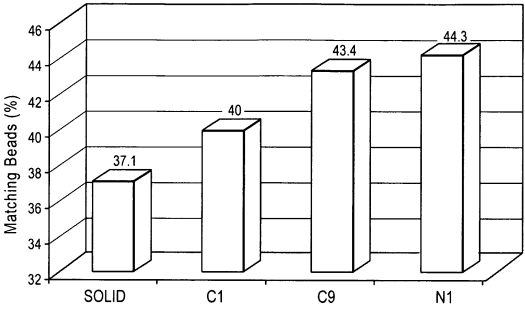


Figure 22

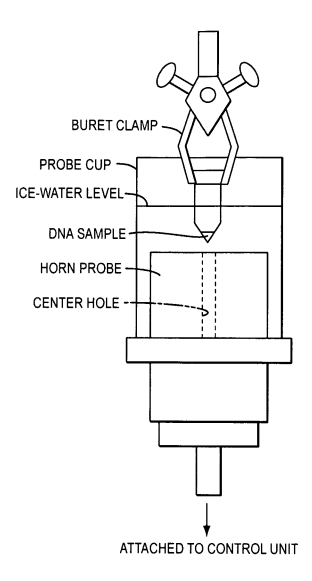
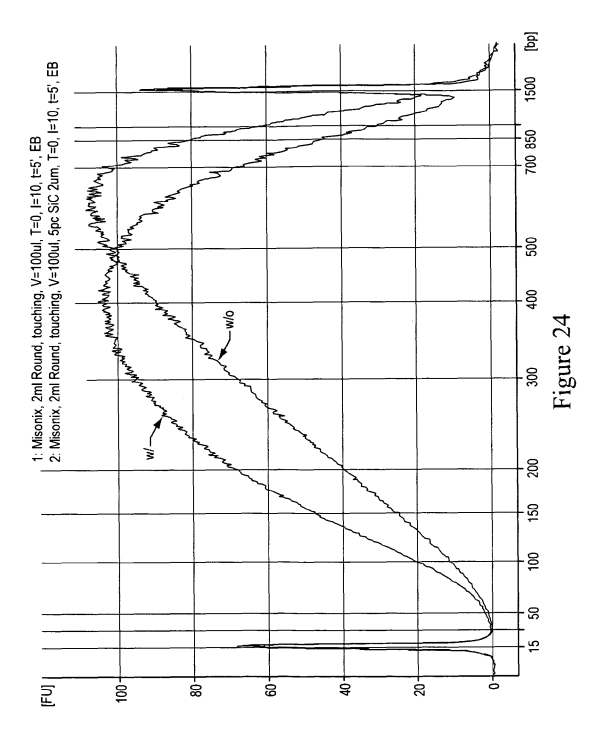
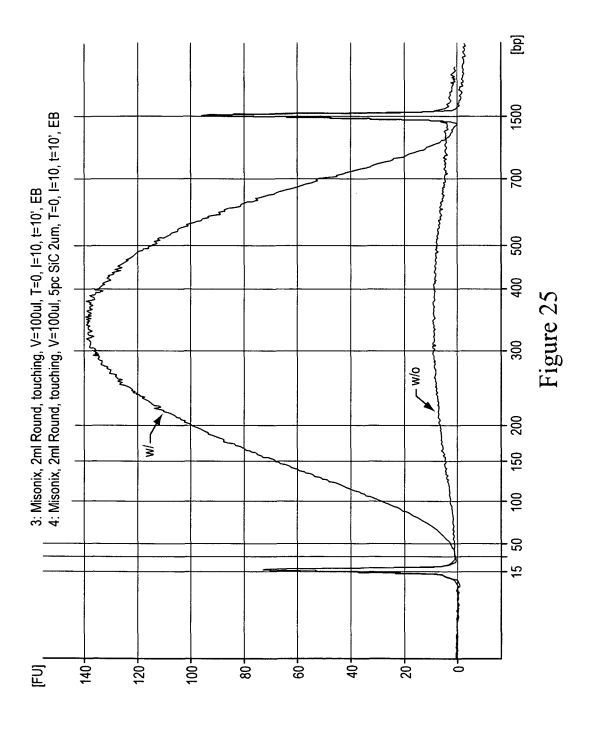
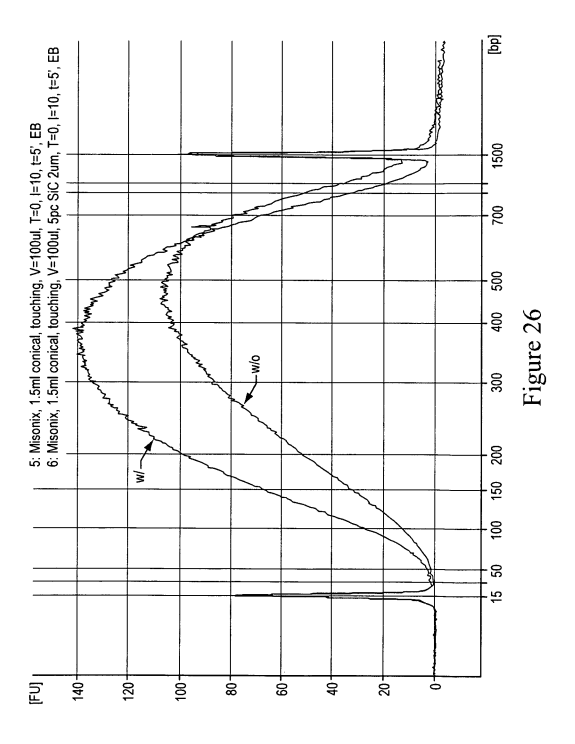
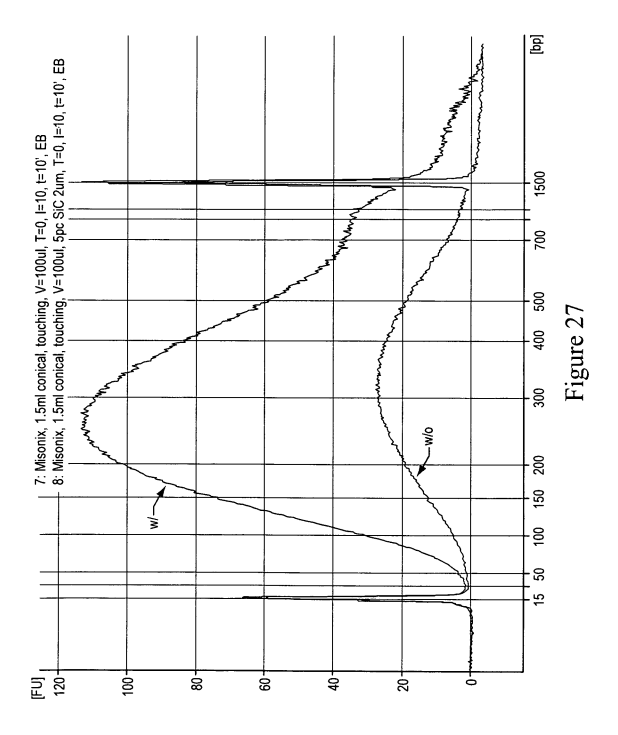


Figure 23









#### METHODS AND APPARATUSES FOR NUCLEIC ACID SHEARING BY SONICATION

#### CROSS-REFERENCE TO RELATED APPLICATIONS

This application claims a priority benefit under 35 U.S.C. §119(e) from U.S. Patent Application No. 61/029,113, filed Feb. 15, 2008, which is incorporated herein by reference.

#### **FIELD**

Methods and apparatuses for nucleic acid fragmentation by sonication, more specifically mechanical DNA fragmentation by sonication with microparticles. The present teachings 13 can be expanded to material for fragmentation that includes any substance including nucleic acid, for example chromatin that includes a nucleic acid and protein, and any long polymer, other than nucleic acid.

#### BACKGROUND

Nucleic acid shearing or fragmentation provides the a first step in several embodiments for constructing nucleic acid libraries, as well as, embodiments for hybridization of target 25 nucleic acids on solid supports, for example microarrays. These embodiments benefit from controlled shearing forces to provide increased efficiency in fragmentation and subsequent analysis. Nucleic acid, for example DNA or RNA, fragmentation are the focus of next-generation sequencing 30 platforms such as those by 454 (Roche Molecular, Inc.), SOLiD (Applied Biosystems), and Solexa (Illumina, Inc.). These platforms each have different embodiments for nucleic acid fragmentation that determine different parameters such as efficiency of fragmentation, fragmentation time, fragment 35 length distribution range, and quality of fragments generated. For example, double-stranded DNA may be accompanied by single-stranded (denatured) DNA, or damaged (depurinated) DNA. Furthermore, there are several applications of nextgeneration sequencing that would benefit from higher 40 throughput that could be achieved by reducing the time needed to process each sample.

There are different methods for nucleic acid fragmentation. Nucleic acids can be fragmented chemically by enzymatic fragmented mechanically, such as by hydrodynamic shearing or sonication. Mechanical fragmentation can occur by several methods known in the art, including shearing of DNA by passing it through the narrow capillary or orifice (Oefner et al., Nucleic Acids Res. 1996; Thorstenson et al., Genome Res. 50 1995), for example a hypodermic needle, sonicating the DNA, such as by ultrasound (Bankier, Methods Mol. Biol. 1993), grinding in cell homogenizers (Rodriguez L V. Arch Biochem Biophys. 1980), for example stirring in a blender, or nebulization. Mechanical fragmentation results, in some 55 methods, in double strand breaks within a DNA molecule. Sonication is used widely for random fragmentation of nucleic acids for sequencing library or microarray probe preparations. A variety of instruments available on the market can provide sonication for nucleic acid preparation. Sonica- 60 tion may also be performed using any convenient approach, e.g., with a multi-tip sonicator or using acoustic sound waves. A Microplate Sonicator® (Misonix Inc.) may be used to partially fragment the DNA. Such a device is described in U.S. Patent Publication No. 2002/0068872. Other examples of sonicators for nucleic acid fragmentation are Vialtweeter or Sonotrode. Another acoustic-based system that may be

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used to fragment DNA is described in U.S. Pat. Nos. 6,719, 449, and 6,948,843 manufactured by Covaris Inc. U.S. Pat. No. 6,235,501 describes a mechanical focusing acoustic sonication method of producing high molecular weight DNA fragments by application of rapidly oscillating reciprocal mechanical energy in the presence of a liquid medium in a closed container, which may be used to mechanically fragment the DNA. An exemplary configuration of such an instrument is shown at FIG. 4 illustrating a concave transducer focusing the acoustic energy through a water bath to a focal zone in the sample liquid contained in the sample vessel. In various embodiments, the focused acoustic energy can have a frequency of 1.1 MHz with 400 Watts of power applied (100 Watt maximum at 20% duty cycle), on a volume of approximately 2 millimeters in diameter by 6 millimeters in height (depth by length). Sonication parameters (such as power, duty cycle, and cycles per burst can be adjusted in the process recipe through software settings. The sonication region shape and volume is hardware dependent and can be modified with hardware changes. Purified nucleic acids can be amplified prior to or after a fragmentation step.

The shearing of a nucleic acid molecule in a liquid medium is achieved through the hydrodynamic action of the liquid on the molecule itself. When a velocity gradient exists within the liquid medium, the shear stresses produced by the elongational components of the flow result into an aligning and extensional action on the nucleic acid molecules along the direction of the shear stresses.

When the applied hydrodynamic action (tensile forces, bending moments, ect) builds up to exceed the intrinsic strength of the polymeric chain, a breakage in the chain will result, giving rise to two fragments, each shorter than the original. In general, since the hydrodynamic action applied to the nucleic acid molecule is proportional to its length, it is increasingly difficult to shear a fragment of a nucleics acid as it becomes shorter and shorter. The shear stress "tau"τ in the fluid giving rise to the hydrodynamic action on the polymeric chain can be expressed according to the following formula:

$$\tau_{xy} = \mu \cdot \left( \frac{\partial u}{\partial y} + \frac{\partial v}{\partial x} \right)$$

digestion, for example, by DNasel. Nucleic acids can be 45 where \u03c4"mu" is the viscosity of the liquid medium and du/dy, and dv/dx, the velocity gradients within the flow field. In order to enhance the shearing action and thus decrease size fragment and increase shearing throughput, the shear stress can be increased by increasing the viscosity of the liquid medium or by increasing the velocity gradients within the flow field. High concentration glycerol solutions are normally used to increase the viscosity of the liquid medium by several orders of magnitude compared to pure water. In addition, mechanisms such as sonication, can be used to produce stronger flow fields than otherwise achievable through more basic mechanical devices. For example, the minimum fragment size obtained in a Hydroshear instrument, where the liquid is force through a small orifice, is of the order of hundreds of base pair. On the other hand, fragments as short as tens of base pairs can be easily obtained with a sonicator thanks to the violence of the flow generated by ultrasonic cavitation. Taking into account the shear being the sum of elongation and rotation to cause stretching and tumbling to produce scission. This results in 5'-CpG-3' preferential cleavage on doublestranded DNA (Grokhovsky, Mol. Bio., 2006). Typical parameters for optimizing nucleic acid fragmentation are sonication parameters (bursts per cycle, intensity, and duty

cycle), process temperature, buffer viscosity, sample volume, nucleic acid amount, sample vessel size and material, buffer ionic strength, and nucleic acid purification method.

Varying lengths of fragments can be provided depending on the sequencing platform. For example, the Illumina 1  $\overset{\circ}{\mathrm{G}}$  5 sequencing platform requires the sonication of pure DNA to generate 100-300 bp pieces for fragment libraries, and of chromatin to get fragments in 200-700 bp range for ChIPsequencing, using i.e. 250 Sonifier (Branson) or Bioruptor (Diagenode AS). Another example, SOLiD used sonication 10 for fragment library preparation to generate size ranges of 60 to 90 base pair fragments from purified nucleic acids. This can be achieved with a Covaris, Inc. S2 sonicator fragmenting the nucleic acid for 40 minutes at maximum setting for power and frequency. Exploiting such instruments at maximum capacity 15 for long periods of time, such as those needed to process each nucleic acid sample, can accelerate instrument aging visible as decline in instrument performance. Further, 40 minute fragmentation cycles (like in the SOLiD protocol) limit the throughput for sequencing instruments by extending the time 20 necessary for gene library generation. In addition, sonication of purified nucleic acids is carried out in glycerol or other viscous liquids to increase the friction on the nucleic acids. However, the glycerol is then separated from the nucleic acid reduce the recovery of the nucleic acid fragments. Furthermore, long fragmentation cycles under high power settings increases the probability and extent of damage and denaturation of the purified nucleic acids (Milowska et al., Biomolecular Engineering 2007). This can be attributed to cavitation 30 induced by sonication. Cavitation can collapse microbubbles, produce microjets, or produce shock waves in the sample liquid, as well as, generating strong flow, localized temperature rise, production of free radicals, for example H and OH (Fuciarelli et al, Free Radical Biology & Medicine, 1995).

It is desirable to provide a method for preparing nucleic acid fragments from a sample of purified nucleic acid that reduces the length of fragmentation time. It is desirable to provide a method for preparing nucleic acid fragments from a sample of purified nucleic acid that avoids glycerol or other 40 viscous liquids and fragments the nucleic acid in an aqueous solution. It is desirable to increase the recovery of fragmented nucleic acids by reducing the power settings of sonicators or reducing the loss of nucleic acid fragments to post-fragmentation separation. It is also desirable to improve sequencing 45 results by reducing the bias of sonication toward certain fragment sizes or fragment types (as opposed to the randomness of nebulization or hydroshearing). The present invention provides these desired results with a method for preparing nucleic acid fragments from a sample of purified nucleic acid 50 by adding particles to the sample and sonicating the suspension. It is counterintuitive that adding particles to the sample would provide the desired fragmentation because the increase in viscosity tends to stop the mechanism for fragmentation at certain point by greatly reducing the molecules spatial turn- 55 over in the focal point of sonication. Therefore, the desirable results of the present invention are not predictable based on current understanding of nucleic acid fragmentation.

Particles have been used with sonication to lyse cells, see for example U.S. Pat. No. 6,440,725 describing a cartridge for 60 cell lysis using beads and an ultrasonic transducer. However, the application of particles in cell lysis via sonication cannot achieve nucleic acid fragmentation because of the cell lysate present. Furthermore, nucleic acid fragmentation is contrary to the goal of cell lysis for detecting low copy of DNA targets 65 in large volume sample, i.e. for diagnostics. To detect lowcopy nucleic acids targets after cell lysis it is desirable to have

the extracted DNA in high molecular weight form, as excessive degradation by i.e. oversonication can dramatically reduce the average DNA fragment length below the length of amplicon, thus greatly reducing the sensitivity of such PCRbased diagnostic methods.

It is also desirable to provide shearing of material that contains some nucleic acid, for example chromatic with DNA and proteins. It is also desirable to provide shearing of other long polymers that are not organic in nature. The present teachings can be expanded to material for fragmentation that includes any substance including nucleic acid, for example chromatin that includes a nucleic acid and protein, and any long polymer, other than nucleic acid where applications require shearing of the polymer for further processing.

#### SUMMARY

In certain embodiments, a method for preparing nucleic acid fragments from a sample of purified nucleic acid is provided. In certain embodiments, a sample of purified nucleic acid is provided, particles are added to the sample, a suspension of the sample and the particles is sonicated, and the nucleic acid fragments are collected.

In certain embodiments, a method for preparing a gene fragments. This process requires chemical extraction and can 25 library from nucleic acid extraction is provided. In certain embodiments, a sample of purified nucleic acid is provided, particles are added to the sample, no nonaqueous solvents are added to the sample, a suspension of the sample and the particles is sonicated, and the nucleic acid fragments are collected.

> In certain embodiments, a method for preparing nucleic acid fragments from a sample of purified nucleic acid is provided. In certain embodiments, a sample of purified nucleic acid is provided which does not include cell lysates, particles are added to the sample, a suspension of the sample and the particles is sonicated, and the nucleic acid fragments are collected.

> In certain embodiments, a kit for preparing nucleic acid fragments from a sample of purified nucleic acid is provided. In certain embodiments, an aqueous buffer solution is included, wherein the solution is substantially free of nonaqueous solvents, and monodisperse particles are included for addition to the buffer solution and the sample of purified nucleic acid.

> The present teachings can be expanded to material for fragmentation that includes any substance including nucleic acid, for example chromatin that includes a nucleic acid and protein, and any long polymer, other than nucleic acid where applications require shearing of the polymer for further processing. These and other features of the present teachings are set forth herein.

#### BRIEF DESCRIPTION OF THE FIGURES

FIG. 1 shows a graph of DNA fragmentation using existing methods of sonication.

FIG. 2 shows a graph of DNA fragmentation comparing existing methods and non-limiting exemplary methods of the present invention.

FIG. 3 shows a non-limiting exemplary protocol for short fragment library preparation.

FIG. 4 shows a non-limiting exemplary apparatus for focusing acoustic sonication.

FIG. 5 shows a graph demonstrating the effects of temperature on fragmentation.

FIG. 6 shows a graph demonstrating the effects of bursts per cycle on fragmentation.

FIG. 7 shows a graph demonstrating the effects of duty cycle on fragmentation.

FIG. 8 shows a graph demonstrating the effects of particle size on fragmentation.

FIGS. 9, 10, and 17 show graphs demonstrating the effects of particle concentration on fragmentation.

FIG. 11 shows a graph demonstrating the effects of particle material on fragmentation.

FIGS. 12, 14, and 19 show graphs demonstrating the effects of time on fragmentation.

FIG. 13 shows a graph demonstrating the effects of viscosity of solvent on fragmentation.

FIGS. **15** through **16** shows graphs demonstrating fragment size distribution and the yield of recovered fragmented DNA depending on fragmentation method used in some <sup>15</sup> embodiments of present invention.

FIG. 18 shows a graph demonstrating the effects of different amounts of purified nucleic acid on fragmentation.

FIGS. **20** and **21** show a graph and a gel image to demonstrate different particle material effects on nucleic acid fragmentation.

FIG. 22 shows the quality of SOLiD sequencing results of fragment libraries constructed using different DNA shearing methods.

FIG. **23** shows a schematic of the cup-and-horn sonicator <sup>25</sup> used to shear DNA and to produce the data in the following figures

FIG. **24** shows a graph of DNA fragmentation comparing existing methods ("w/o") and non-limiting exemplary methods ("w") of the present invention. The fragmentation was <sup>30</sup> performed with a 5' run in a 2 ml round bottom Eppendorf tube in a Misonix cup-and-horn sonicator.

FIG. **25** shows a graph of DNA fragmentation comparing existing methods ("w/o") and non-limiting exemplary methods ("w") of the present invention. The fragmentation was <sup>35</sup> performed with a 10′ run in a 2 ml round bottom Eppendorf tube in a Misonix cup-and-horn sonicator.

FIG. **26** shows a graph of DNA fragmentation comparing existing methods ("w/o") and non-limiting exemplary methods ("w") of the present invention. The fragmentation was 40 performed with a 5' run in a 1.5 ml conical bottom Eppendorf tube in a Misonix cup-and-horn sonicator.

FIG. 27 shows a graph of DNA fragmentation comparing existing methods ("w/o") and non-limiting exemplary methods ("w") of the present invention. The fragmentation was 45 performed with a 10' run in a 1.5 ml conical bottom Eppendorf tube in a Misonix cup-and-horn sonicator.

## DETAILED DESCRIPTION OF CERTAIN EMBODIMENTS

The section headings used herein are for organizational purposes only and are not to be construed as limiting the subject matter described. All documents, or portions of documents, cited herein, including but not limited to patents, 55 patent applications, articles, books, and treatises, are hereby expressly incorporated by reference in their entirety for any purpose. In the event that one or more of the incorporated documents or portions of documents defines a term that contradicts that term's definition in this application, this application controls.

The use of the singular includes the plural unless specifically stated otherwise. The word "a" or "an" means "at least one" unless specifically stated otherwise. The use of "or" means "and/or" unless stated otherwise. The meaning of the 65 phrase "at least one" is equivalent to the meaning of the phrase "one or more." Furthermore, the use of the term

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"including," as well as other forms, such as "includes" and "included," is not limiting. Also, terms such as "element" or "component" encompass both elements or components comprising one unit and elements or components that comprise more than one unit unless specifically stated otherwise.

According to various embodiments, FIG. 3 illustrates an exemplary protocol for DNA fragment library preparation, for example SOLiD sequencing platform. The process shown takes genomic DNA that has been purified to remove any cellular material that resulted from lysis to extract the genomic DNA. The DNA is then sheared to provide fragments. The typical shearing protocol includes taking a 500 microliter sample that contains 10 micrograms of DNA in an 85% glycerol solution, and using a Covaris sonicator to generate roughly 100 watts of energy for a 40 minutes cycle. The DNA is sheared and then the ends are repaired and ligated with adaptors and undergo nick translation to create P1-P2 ligated library molecules. Nick translation removes any nicks left in the dsDNA molecules during upstream steps of library construction in order to allow for efficient final PCR amplification of the library (see Large scale PCR step in FIG. 3). The library containing a mixture of molecules of various sizes is then resolved upon 4% agarose gel electrophoresis and molecules in 120-160 base pair range are size selected and extracted from agarose. The library can be subjected to trial and large-scale PCR and then further purification. The library is then taken to quality control by for example electrophoretic gel or a Bioanalyzer (Agilent Technologies). According to various embodiments, a method of forming a released paired tag from a DNA fragment of interest is provided. The method comprises ligating a first adaptor to a first end of a DNA fragment of interest and ligating a second adaptor to a second end of the DNA fragment of interest, thereby producing an adaptor modified fragment, circularizing the adaptor modified fragment by attaching a third adaptor to the adaptor modified fragment, thereby forming a circular nucleic acid molecule wherein a first nick is present between the third adaptor and the first adaptor and a second nick is present between the second adaptor and the third adaptor, wherein the circular nucleic acid molecule comprises a first strand and a second strand of DNA, and wherein the first nick and the second nick are not present on the same strand of the circular nucleic acid molecule; performing a nick translation reaction wherein a nick on each strand of the circular nucleic acid molecule is translated into the DNA fragment of interest, and cleaving the circular nucleic acid molecule at a position at a translated nick, thereby forming a released paired tag. Additional examples are contained in copending application titled, "METHOD OF MAKING A PAIRED TAG LIBRARY FOR 50 NUCLEIC ACID SEQUENCING" with Ser. No. 12/350,837, filed Jan. 8, 2009.

TE buffer is a commonly used buffer solution in molecular biology, especially in procedures involving DNA or RNA. It is called "TE" buffer because it contains Tris, a common pH buffer, and EDTA, a molecule chelating cations like Mg2+. The purpose of TE buffer is to protect DNA or RNA from degradation. A typical recipe for making TE buffer is:10 mM Tris, bring to pH 7.5 with HCl, 1 mM EDTA. Based on nuclease studies from the 80's, the pH is usually adjusted to 7.5 for RNA and 8.0 for DNA. The respective DNA and RNA nucleases are supposed to be less active at these pH values. But pH 8.0 can safely be used for storage of both DNA and RNA. EDTA further inactivates nucleases, by binding to metal ions required by these enzymes.

As used herein, the terms "nucleic acid" means singlestranded and double-stranded polymers of nucleotide mono-

mers, including, but not limited to, 2'-deoxyribonucleotides and ribonucleotides (RNA) linked by internucleotide phosphodiester bond linkages, e.g. 3'-5' and 2'-5', inverted linkages, e.g. 3'-3' and 5'-5', branched structures, or analog nucleic acids. Nucleic acids can be natural nucleic acids, 5 artificial nucleic acids, analogs thereof, or combinations thereof. Nucleic acids have associated counter ions, such as H<sub>+</sub>, NH4<sub>+</sub>, trialkylammonium, Mg2<sub>+</sub>, Na<sub>+</sub> and the like. A nucleic acid can be composed entirely of deoxyribonucleotides, entirely of ribonucleotides, or chimeric mixtures thereof. Nucleic acid can be comprised of nucleobase and sugar analogs. Nucleic acid typically range in size from a few monomeric units, e.g. 5-40 when they are more commonly frequently referred to in the art as oligonucleotides, to several thousands of monomeric nucleotide units. Unless denoted 15 otherwise, whenever a nucleic acid sequence is represented, it will be understood that the nucleotides are in 5' to 3' order from left to right and that "A" denotes deoxyadenosine, "C" denotes deoxycytidine, "G" denotes deoxyguanosine, and "T" denotes thymidine.

Nucleic acids are said to have "5' ends" and "3' ends" because mononucleotides are reacted to make nucleic acids in a manner such that the 5' phosphate of one mononucleotide pentose ring is attached to the 3' oxygen of its neighbor in one nucleic acid is referred to as the "5' end" if its 5' phosphate is not linked to the 3' oxygen of a mononucleotide pentose ring and as the "3' end" if its 3' oxygen is not linked to a 5' phosphate of a subsequent mononucleotide pentose ring. As used herein, a nucleic acid sequence, even if internal to a 30 larger nucleic acid, also can be said to have 5' and 3' ends.

As used herein, "purified nucleic acid" denotes a genomic polynucleotide without cellular material. For example, a sample from a cell, where the polynucleotide is fragmented into acceptable sizes to serve as nucleic acid fragments. The 35 options and variations of purification of the genomic polynucleotide are broadly known to one skilled in the art of cellular lysis and vary on the cellular material and inhibitors that can be contained in that material.

As used herein, the term "chromatin" encompasses com- 40 plex of DNA and protein that makes up chromosomes. It is found inside the nuclei of eukaryotic cells, and within the nucleoid in prokaryotes. The major proteins involved in chromatin are histone proteins, although many other chromosomal proteins have prominent roles too. The term also 45 encompasses DNA/protein/RNA complex extracted from eukaryotic lysed interphase nuclei. Just which of the multitudinous substances present in a nucleus will constitute a part of the extracted material will depend in part on the technique each researcher uses. Furthermore, the composition and properties of chromatin vary from one cell type to the another, during development of a specific cell type, and at different stages in the cell cycle. The term also encompasses DNA double helix in the cell nucleus is packaged by special proteins termed histones. The formed protein/DNA complex is 55 called chromatin. The structural entity of chromatin is the

Chromatin is DNA plus the proteins (and RNA) that package DNA within the cell nucleus.

As used herein, the term "microarray" encompasses an 60 arrangement of polynucleotides present on a solid support or in an arrangement of vessels. Certain array formats are referred to as a "chip" or "biochip" (M. Schena, Ed. Microarray Biochip Technology, BioTechnique Books, Eaton Publishing, Natick, Mass. (2000)). An array can comprise a low- 65 density number of addressable locations, e.g. 1 to about 12, medium-density, e.g. about a hundred or more locations, or a

high-density number, e.g. a thousand or more. Typically, the array format is a geometrically-regular shape that allows for fabrication, handling, placement, stacking, reagent introduction, detection, and storage. The array can be configured in a row and column format, with regular spacing between each location. Alternatively, the locations can be bundled, mixed, or homogeneously blended for equalized treatment and/or sampling. An array can comprise a plurality of addressable locations configured so that each location is spatially addressable for high-throughput handling, robotic delivery, masking, and/or sampling of reagents and/or by detection means including scanning by laser illumination and confocal and/or deflective light gathering. The array can comprise one or more "addressable locations," e.g., "addressable positions," that is, physical locations that comprise a known type of molecule.

#### Certain Exemplary Embodiments

In each of the following embodiments, markers were used for 15 base pairs and 1500 base pairs to demonstrate the relationship between time in seconds on the x-axis and the length of nucleic acid fragments in base pairs. These will be reflected as sharp spikes at the ends of each graph.

Certain Exemplary Embodiments of Sonication

In various embodiments according to the present invention, direction via a phosphodiester linkage. Therefore, an end of a 25 a method for nucleic acid fragmentation is provided which includes adding particles to the sample of purified nucleic acid. The present invention has demonstrated accelerated fragmentation times of at least 8 fold to at least 16 fold. For example, the standard 40 minutes sonication was significantly shortened. The analysis for fragmentation, according to various embodiments, was measured using DNA 100 Assay chips and reagents (Agilent) on a Bioanalyzer (Agilent). FIG. 1 illustrates nucleic acid fragmentation in 85% glycerol with parameters of 20% duty cycle, intensity of 10%, cycle of 60 seconds, and temperature of 18 degrees Celcius. The curve (I) shows the size distribution of fragments generated after a standard 40 minute procedure used in SOLiD at the highest acoustic frequency of 1000 bursts per cycle. The high molecular weight (for example greater than 15,000 bases) genomic DNA was converted to short fragments (peak at 48 base pairs). The curve (III) shows the same conditions except for reducing the time to 5 minutes. The fragmented nucleic acid is spread between the 15 base pair and 1500 based pair markers with a majority remaining in the high size range (peak at approximately 500 base pairs). The curve (II) shows an embodiment decreasing the frequency to 50 bursts per cycle that shifted the fragment distribution towards the low range compared to curve (III) but still not providing a majority of DNA fragments in the desired range of 60 base pairs to 90 base pairs, for example in preparing a fragment library for SOLiD. Curve (II) is also shown on FIG. 2 as curve (IV).

In various embodiments, the following standard SOLiD protocol test conditions were followed: DNA amount of 10 micrograms, E. coli strain Novablue (Novagen) DNA as sample nucleic acid, phenol-chloroform DNA purification method, 500 microliter sample volume, 85% glycerol, 1.6 micrometer monodispersed glass particles (Duke Scientific) vial sample vessel with dimensions of 65 millimeters by 13 millimeters. The sonication parameters were 40 minutes, at 20% duty cycle with intensity of 10, and 1000 bursts per cycle. FIG. 5 illustrates a graph showing temperature in degrees Celsius and time in minutes. This embodiment demonstrates that even if sample temperature were to vary, time is the dominant parameter. The parameter of 18 degrees Celsius was selected for other exemplary embodiments. FIG. 6 illustrates a graph showing the effect of frequency (bursts per

cycle) and time in minutes on shearing of DNA. This embodiment demonstrates that lower bursts per cycle provide the highest yield for the desired fragmentation at 60 base pairs to 90 base pairs. The parameter of 50 bursts per cycle was selected for other exemplary embodiments. FIG. 7 illustrates 5 a graph showing the effect of duty cycle on fragmentation of nucleic acids. This embodiment demonstrates that duty cycle affects fragmentation. The parameter of 20% was selected for other exemplary embodiments.

In various embodiments, a Covaris S2 sonicator was used 10 with glass tubes immersed in a water bath at 18 degrees Celcius. The sample included 500 microliters of 1×TE buffer pH 8.0, 10 micrograms of genomic DNA isolated from E. coli strain NovaBlue (Novagen), 85% glycerol, 1.6 micrometer monodispersed glass particles (Duke Scientific) were added 15 to the sample (as dry powder) to provide a 5% (weight/ volume) suspension to the embodiment of curve (IV) (same as curve (II) in FIG. 1) and fragmented the nucleic acid as shown in curve (V) of FIG. 2. Further time reduction to 2.5 minutes, as shown by curve (VI) was achieved without sub- 20 stantially changing the fragment size distribution shown in curve (V). In various embodiments, FIG. 12 illustrates a graph showing the minimization of time had on fragmentation. This embodiment demonstrated that duration for fragmentation shorter than 2 minutes decreased the fragmentation 25 at the desired fragment sizes of 60 base pairs to 90 base pairs.

In various embodiments, the size of the particles can range from 0.1 micrometers to 100 micrometers and provide a concentration of 0.1% to 20% weight per volume during fragmentation. In various embodiments, FIG. 8 illustrates a graph 30 showing the effect of particle size at a fixed concentration of 2%. This embodiment demonstrates that lower particle sizes, e.g. beads 1.6 micrometers to 8 micrometers of diameter, provided the desirable fragment sizes of 60 base pairs to 90 base pairs. In various embodiments, FIGS. 9 and 10 illustrate 35 graphs showing the effect particle concentration at a fixed particle size of 1.6 micrometers at 5 degrees Celsius and 18 degrees Celsius, respectively. This embodiment demonstrated that particle concentration 2% to 5% provided the desirable fragment sizes. In various embodiments, FIG. 19 40 illustrates the dependence on time for fragmentation with 2.0 micrometer glass particles in glycerol-free solution.

In various embodiments, the particles can be spherical or irregular shaped, non-porous or partial porosity, monodispersed or broadly dispersed over a size range. In various 45 embodiments the composition of the particles can include glass (e.g. borosilicate glass), silicon carbide, silica, soda lime glass, diamond, metal (e.g. Ag, Au, Al, Co, Cu, Fe, Mo, Ni, Ti, W, Zn), metal oxides (e.g. Al<sub>2</sub>O<sub>3</sub>, ZrO<sub>2</sub>, HfO<sub>2</sub>, TiO<sub>2</sub>, CeO<sub>2</sub>, SiO<sub>2</sub> and mixed oxides), hollow ceramic, metal (e.g. Al 50 or Ni) coated hollow ceramic or combinations thereof. In various embodiments, the particles can be non-magnetic or paramagnetic. In various embodiments, FIG. 11 illustrates a graph showing the effect of particle composition on fragmentation. This embodiment demonstrated that silica, even at 55 particle sizes that had provided favorable fragmentation, did not fragment nucleic acids at the desired fragment sizes as did 1.6 micrometer glass particles. In various embodiments, FIGS. 20 and 21 illustrate silicon carbide and borosilicate glass as alternative compositions for the particles. Silicon 60 carbide was obtained from Electro Abrasives Co. with a hardness of 9.6 (Moh) and a blocky, sharp shape and borosilicate glass from Duke Scientific Inc. with a hardness of 6.5 (Moh) and a spherical shape. FIG. 21 shows a gel electrophoresis showing the fragment size for the nucleic acid.

In various embodiments, it is desirable to reduce the concentration of glycerol to minimize losses of DNA during 10

subsequent step of DNA-binding column purification, i.e. QIAGEN MinElute column, prior to end-repair of sonicated DNA provided to for example, sequencing analysis by, for example, SOLiD. Even more desirable is to fragment DNA in glycerol-free solution in order to couple the sheared DNA sample directly to the end-repair enzymatic reaction, thus, eliminating without the intermediate column step, which would further increase the yield of fragmented nucleic acid provided to for example, sequencing analysis by, for example, SOLiD. In various embodiments, FIG. 13 illustrates a graph showing the effect of glycerol concentration on fragmentation in the presence of glass beads. The sonication time was fixed for each sample to 2.5 minutes. This embodiment demonstrates that some concentration of glycerol particularly 60% to provide an effect (increased yield in the target range) at that time frame. In various embodiments, FIG. 14 illustrates a graph showing the effect of time at a fixed glycerol concentration of 60%. This embodiment demonstrates that increasing the sonication time provides fragments closer to the desirable fragment sizes. The following chart shows several embodiments for fragmentation by sonication with different concentrations of glycerol as compared to nebuliza-

Treatment	Method	Time	Glycerol	Beads
C1	Covaris	2'	60%	5%* 1.6 um (Duke#145)
C9	Covaris	30'	0%	5%* 1.6 um (Duke#145)
N1	Nebulizer (75 psi)	15'	75%	None
Solid	Covaris	40'	85%	None

In various embodiments, FIG. 15 illustrates a graph with nucleic acid fragment size distribution generated under conditions shown in the above chart. This embodiment demonstrates that sonication without glycerol or less glycerol than 85% provides better yields of nucleic acid fragments in the desired size range for subsequent analysis. FIG. 167 illustrates a comparison of yield of total and double stranded DNA ethanol precipitated after fragmentation by different methods. UV absorbance measures total yield of nucleic acid (single stranded DNA, double stranded DNA, and free nucleotides combined), while, PicoGreen is highly selective for double stranded DNA. Only double-stranded DNA generated during fragmentation step will serve as a substrate for further steps of library construction. As evident from the FIG. 16 the highest yield of total and double stranded DNA is achieved with method C9. In contrast, the standard SOLiD method results in significant losses due to overshearing and thermal denaturation of DNA—only 8.7% of input DNA is recovered with 17% of that in double stranded form. The following chart illustrates a comparison for quantity of double stranded DNA in a target size range captured by the different embodiments.

Treatment	Time	Glycerol	Beads	60-90 bp ds-DNA (ng)
C1	2'	60%	5% 1.6 um	196
C9	30'	0%	5% 1.6 um	785
N1	15'	75%	None	254
V (Solid)	40'	85%	None	15.5

The sequencing data captured shown in the FIG. 22 shows the improved yield of Matching Beads (as percentage of Readable Beads) in C9 method compared to SOLiD method, demonstrating that the sequencing data of better quality is obtained when using the combination of glycerol-free solu-

tion with microbeads. In various embodiments, the graph in FIG. 17 shows an effect of bead concentration on fragmentation of the nucleic acid in an aqueous solution with out non-aqueous solvent such as glycerol.

FIG. 23 shows a schematic representation of ing a cup and 5 horn sonicator. A sonicator of the cup-and-horn type is traditionally used to shear DNA down to sizes of ~500 bp. It is well known that the amount of ds-DNA and its integrity dramatically decreasesdecrease if this method is used to obtain shorter fragments. (CSH protocols, 2006).

FIG. 24 shows the distribution of DNA fragment sizes obtained after shearing DNA for 5 minutes' at continuous maximum intensity in a Misonix 3000 cup and horn sonicator. The curve marked "w" was obtained with an embodiment consisting of 5% Silicon Carbide beads of 2 um average size 15 obtained from Electroabrasives and suspended in 200 ul of aqueous buffer placed in a 2 ml round bottom Eppendorf tube, while the one marked "w/o" was obtained using the traditional method, subjecting the sample to the same conditions but without beads. This figure demonstrates that fragments 20 30% shorter at the distribution peak can be achieved with this embodiment (400 vs 600 bp).

FIG. 25 shows the distribution of DNA fragment sizes obtained after shearing DNA for 10 minutes at continuous maximum intensity in a Misonix 3000 cup and horn sonicator. 25 The curve marked "w" was obtained with an embodiment consisting of 5% Silicon Carbide beads of 2 um average size obtained from Electroabrasives and suspended in 200 ul of aqueous buffer placed in a 2 ml round bottom Eppendorf tube, while the one marked "w/o" was obtained using the tradi- 30 tional method, subjecting the sample to the same conditions but without beads. This figure demonstrates that without the embodiment claimed in this invention the DNA greatly denatures and only a small fraction of the input material survives sonication. On the other hand, when the non-limiting 35 described embodiment is used, the amount of sheared DNA in double stranded form is preserved, thus providing a large yield. In general, to prevent DNA from denaturing, the ultrasonic energy is applied intermittently in pulses to allow the liquid to cool down in between pulses, thus mitigating the 40 thermally induced DNA denaturing process. This translates into a longer processing time: for example, if a duty cycle of 0.5 is used, the total processing time will be twice as long. A clear benefit of this non-limiting embodiment is that sonication can be applied at continuous maximum power to the 45 sample without it undergoing measurable degradation.

FIG. 26 shows the distribution of DNA fragment sizes obtained after shearing DNA for 5 minutes at continuous maximum intensity in a Misonix 3000 cup and horn sonicator. The curve marked "w" was obtained with an embodiment 50 consisting of 5% Silicon Carbide beads of 2 um average size obtained from Electroabrasives and suspended in 200 ul of aqueous buffer placed in a 1.5 ml conical bottom Eppendorf tube, while the one marked "w/o" was obtained using the traditional method, subjecting the sample to the same conditions but without beads. This figure demonstrates that fragments 30% shorter at the distribution peak can be achieved with this embodiment (400 vs 600 bp).

FIG. 27 shows the distribution of DNA fragment sizes obtained after shearing DNA for 10 minutes at continuous 60 maximum intensity in a Misonix 3000 cup and horn sonicator. The curve marked "w" was obtained with an embodiment consisting of 5% Silicon Carbide beads of 2 um average size obtained from Electroabrasives and suspended in 200 ul of

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aqueous buffer placed in a 1.5 ml conical bottom Eppendorf tube, while the one marked "w/o" was obtained using the traditional method, subjecting the sample to the same conditions but without beads. This figure demonstrates that without 5 the embodiment claimed in this invention the DNA greatly denatures and only a small fraction of the input material survives sonication. On the other hand, when the non-limiting described embodiment is used, the amount of sheared DNA in double stranded form is preserved, thus providing a large 10 yield.

What is claimed:

1. A method for preparing nucleic acid fragments from a sample of purified nucleic acid, the method comprising:

providing a sample of purified nucleic acid;

adding particles to the sample;

sonicating a suspension of the sample and the particles to produce nucleic acids fragments; and

collecting at least a portion of the nucleic acid fragments.

- 2. The method of claim 1, wherein the particles have a size from about 0.1 micrometers to about 100 micrometers.
- 3. The method of claim 1, wherein the particles are mono-dispersed.
- 4. The method of claim 1, wherein the particles comprise at least one of glass, borosilicate glass, silica, soda lime glass, carbamide, metal, metal oxide, ceramic, and metal-coated ceramic.
- 5. The method of claim 4, wherein the particles are non-magnetic or paramagnetic.
- 6. The method of claim 1, wherein the sonicating comprises focusing acoustic sonication.
- 7. The method of claim 1, wherein the suspension is aque-
- **8**. The method of claim **1**, wherein the suspension comprises less than 85% glycerol.
- 9. The method of claim 1, sonicating the suspension is less than 5 minutes to provide from about 60 to about 90 base pair fragments.
- 10. The method of claim 1, further comprising sequencing the nucleic acid fragments.
- 11. The method of claim 1, further comprising hybridizing the nucleic acid fragments on a microarray.
- 12. The methods of claims 1, wherein sonication comprises subjecting the suspension to an ultrasonic field.
- 13. The methods of claims 1, wherein the fragment sizerange is controlled by particle size, particle chemical composition, percentage of particles in suspension, viscosity of solution, time of treatment, and sonication parameters.
- **14**. A method for preparing nucleic acid fragments from a sample of purified nucleic acid, the method comprising:

providing a sample of purified nucleic acid, wherein the sample does not include cell lysates;

adding particles to the sample;

sonicating a suspension of the sample and the particles to produce nucleic acids fragments; and

collecting at least a portion of the nucleic acid fragments.

15. A method for preparing polymer fragments, the method comprising:

providing a sample of polymer;

adding particles to the sample;

sonicating a suspension of the sample and the particles to produce polymer fragments; and

collecting at least a portion of the polymer fragments.

\* \* \* \* \*

# UNITED STATES PATENT AND TRADEMARK OFFICE CERTIFICATE OF CORRECTION

PATENT NO. : 9,127,306 B2

APPLICATION NO. : 12/372464

DATED : September 8, 2015

INVENTOR(S) : Bashkirov

It is certified that error appears in the above-identified patent and that said Letters Patent is hereby corrected as shown below:

In the claims

Column 12, line 43, Claim 12, delete "s" from "methods"; and delete "s" from "claims" and should read as follows:

12. The method of claim 1, wherein sonication comprises subjecting the suspension to an ultrasonic field.

Column 12, line 45, Claim 13, delete "s" from "methods"; and delete "s" from "claims" and should read as follows:

13. The method of claim 1, wherein the fragment sizerange is controlled by particle size, particle chemical composition, percentage of particles in suspension, viscosity of solution, time of treatment, and sonication parameters.

Signed and Sealed this Eleventh Day of October, 2016

Michelle K. Lee

Michelle K. Lee

Director of the United States Patent and Trademark Office